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# The NPI-1/NPI-3 (Karyopherin α) Binding Site on the Influenza A Virus Nucleoprotein NP Is a Nonconventional Nuclear Localization Signal

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Two cellular proteins, NPI-1 and NPI-3, were previously identified through their interaction with the influenza virus nucleoprotein (NP) by using the yeast two-hybrid system. These proteins were then shown to act as general transport factors (karyopherin α) and nuclear pore-docking proteins to facilitate the transport of the NP and of viral RNA into the nucleus. The yeast two-hybrid assay has now been used to identify the specific domains on the NP that bind to the NPI proteins. Mutational analysis including alanine scanning identified the motifs SxGTKRSYxxM and TKRSxxxM, which are required for binding to NPI-1 and NPI-3, respectively. These sequences were shown to possess nuclear localization signal (NLS) activity following expression of fusion proteins in HeLa cells. These sequences represent a novel nonconventional NLS motif. Another NLS activity not mediated by the NPI binding sites is associated with noncontiguous sequences in the NPI.

The influenza A virus has a segmented genome of eight negative-strand RNAs which are packaged into virions as ribonucleoproteins (RNPs). In addition to RNA, RNP contains the viral nucleoprotein (NP) and the three subunits of the RNA-dependent RNA polymerase, PB1, PB2, and PA. When the influenza virus infects cells, it must ultimately deliver its genetic information to the nucleus, where viral RNA transcription and replication take place. In an attempt to learn more about intracellular host-virus interactions which are required for virus replication and which might affect virulence characteristics, we used the yeast two-hybrid system to search for cellular proteins that interact with the viral NP (20, 21). Two NP-interacting proteins (NPI-1 and NPI-3) were identified and shown to facilitate the nuclear import of the viral NP and-in conjunction with the NP-of viral RNA by using digitoninpermeabilized cells. Independent isolation and characterization of NPI-1 and NPI-3 from mammalian cells and of homologs in Xenopus and Drosophila (3, 4, 10, 15, 18, 27) revealed that these two proteins are general transport factors belonging to the karyopherin a/importin 60 protein family and are involved in the nuclear import of nuclear localization signal (NLS)-containing proteins. In this study we mapped the binding site of NPI-1 karyopherin of hSRP-1 (3, 18, 20, 27) and of NPI-3 karyopherin  $\alpha 2$  Reh-1 (4, 18, 21) on the influenza A virus NP. Mutational analysis of the NP by the yeast two-hybrid system and expression of NP mutants in mammalian cells helped to define an unusual motif in the N terminus of the viral NP which has NLS activity.

# MATERIALS AND METHODS

Yeast, bacterial strain, and plasmids. Sus-hanomyces (cocrisio) PGY4s (M416) np1 ma Ons 3414 CPJ 1X-top644442, plasmids pEG202, p8418-33, and pRFHML, and the HeL a cell cDNX library constructed in pEG4.5 kindly provided by R. Brent (Harvied Medical Schood) have been used previously (20) NP1 Lind NP13 cDNAs cloned in pEG4.5 were isolated from a HeL a cell cDNA.

library (20). The Excherichia volts train used for cloning was DH5a [F=480dlac2, 2M15 2dlac2YA argh]t 169 deak recAl and Al hidRE(r, m, 1) supl. 44 x - the Lgo 496 rel H] (GIBCO BR1)

Construction of recombinant DNA molecules, NP deletion mutants shown in Fig. Ia with constructed between the FeoRI and Viol restriction sites of pEG202 a Tolloys, for my ant A, pBS NP, which contains the influence A PR 8.34 virus 5.0 gove in pldue crip 5K+ (Stratagene Cloring Systems), was cut by BifH (at the leotide positions 80) and 1061) and religited. The deleted NP was then subclosed into pl G202, for mutant B, pBS-5-P was cut by Bg/H and HadIII at usel estide positions \$36 and 1114, respectively, blunted by the Elenow tragment (1L) oili DNA John (rase L) and religated. The deleted NP was then subcloned into \$1.6202. For metants C. F. and F. tragments of the "A" with the b cakpours indicated below were subcloned as PCR products from pBS-NP following emphrication with the appropriate object relectides. Murant I was cloudd is a Bgill Ahol restriction frigorian into the BamHI and Ahol restriction sites of p1 G202. For mutart D. p186-NP was digested with PpuMI. Flurned with Klenow, for I their digested with LeaR1, p1 G202 was digested with Vlot, blunted with LeaR1, p1 G202 was digested with Vlot, blunted in the control of the control with Kler (v. and then digested with Leaf-I. The NP tragment was then sub-Canadam opt G202. For capte siot in manimalian cells (Fig. 1b), the EcoRI to Who I restriction fragments containing the wild type NP and deletion narrants A. B. and by were subcloned into pcDSAI Amp (Invitis gen Corp.).

NP deletion mutants used for Fig. 2 were constructed by PCF between the restriction, actes indicated below. For construction of the fusion generated in the yeast two tybord assay, the PCF products of mutants G. H. J. K. L. and M were cut by B. II Moof and the PCF product of mutant 5, was cut by B mill Moof. These r., tection fragments were closed more the RamHI Moof acts o. pl. G.202. For constructions used in in non-aborescence assays, the PCF products of mutants G. H. J. K. L. and M were cut by BgHLS. L and the PCF products of mutants G. M. L. K. L. and M were cut by BgHLS. L and the PCF products of mutants G. M. and Moof acts of the PCF products of mutants G. M. and the pcf. and the pcf. product of mutants G. M. and the pcf. L and Moof point of the BgHLS. If and the PCF product of mutants G. M. and G. T. L. L. L. L. and M. and all was expression of the P. Inaginents as president on PCF fusion products placed by the pcf. D. R. L. and M. A. L. L. A. L. L. A. L. A

Then is them and deletions in the Past To annow seeds of the NP. In win in Fig. and 4. The class constructed by PCF. For the years, we have I is another PCR product very cut by B/H Mod, and closed into the Bantiti Mod, site, of ITG20. Testitute in pLocation in type III based genes. For the immonathiorises, the assay the PCF products were unity by III Ball and closed into the BgH Sall safes of PLC1. (2) These constructs allowed the expression of the NP tragments as includes of the Putsion production in ILAT recills. All the constructs were continued by fined onder sequence a taky is

Mat. its D\*2 and DF2 vwere generated by PCR it on pBS SP and mutant A respect tely. Mutants D\*2, DF2 v. and O\*3 or t. clery, damap I G202 for use in the versitive behind system and men pcDS M. Anal for expression in manimum and cells Mutant O\*3 was also clouded as a rice is essential. P protein bestoa in ecclor PFCI

Interaction of NP mutants and NPI-1 or NPI-3. The interaction between NP motant and NPI-1 or NPI-3 are assured as described previously (20, 21) briefly, strain I GYBs was transformed with fusion zero, plasmads in addition to the proof a transformed plasmad pSHI-8-34 by a high efficiency lithium acceptable process.

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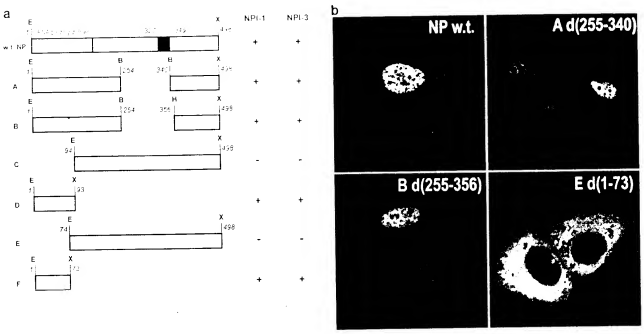


FIG. 1. Amino acids 1 to 73 of the influenza A virus NP contain the binding site for NPI-1 and NPI-3. (a) Interactions of NP deletion mutants and NPI-1 and NPI-3 proteins in the yeast two-hybrid system. Yeast strain EGY48 was cottansformed with pEG202 (containing the different LevA-NP tusion genes), the reporter plasmid pSH18s-34, and pHB34 (containing the NPI-1 cDNA in pIG4-5) or pHB81 (containing the SPI-3 cDNA in pIG4-5) (20). The lengths of the NP fragments are indicated by animo acid positions in the wild-type (w.t.) NP. Binding activity was determined by cerecting the induction of Begalactosidase. A plus sign indicates that a strong blue cofor developed within 12 h following streaking on N-Cral-containing platers, a minus againdicities no reaction. The shaded box indicates RNA binding domains (Lt. 14). The black box indicates a putative NLS (5). Restriction enzyme sites used for subcloning of NP fragments are indicated as follows: B. BgHt. E. EcoRt, H. HindIII. and N. Xhot. (b) Localization of NP deletion mutants in HeLa cells were transfected with pcDNAL Amp, which contains NP deletion mutants and analyzed by indirect immunofluorescence issay 24 h posttransfection, using anti-NP polyclonal antibody for construct L d(1-73) or monoclonal antibody wild-type 'NP and construct SA and B.

tate protocol (13) and plated on glocose-free withictic dropout plates lacking historine, oracid, and tryptophan. In order to away for β galactosidase activity the transformed yeast cells were isolated 2 days posttransformation and streaked onto galactose plates tacking historine, in (cl. and tryptophan and containing 5-bromos-4 chloro-3-indolyl-β-b-galactoside. (N-Gal) (2). The NP mutanic (cloned as LexA fusion genes) that induced β-galactosidase only in the presence of galactose in conjunction with NPI-1 or NPI-3 (cloned as lusion genes with a transcriptional activation domain) were considered to retain the ability to interact with NPI-1 or NPI-3.

Localization of expressed proteins by indirect immunofluorescence assay. HeLa cells were grown on glass coverships to 50 to 85% confluency and transfected with the different constructs (described above) by using Lipotectin according to the manufacturer's recommendations (GIBCO BRL). Twenty-four hours after transfection, the cells were fixed and perincabalized for 15 min in acctone-methanol (1.1) at ±20 °C. The cells were then treated with polyclonal or monoclonal antibodies. Purified monoclonal are body PT107 (unpublished data) or polyclonal rabbit anti-X-31 antibody (23) was used for staming of the NP. The epitope on the NP, which is bound by PT107, has not been mapped. Rabbit anti-SQL PV1 against the measles virus P protein was a kind gift of Dabus 1. Briedis (McGiff University, Montreal, Canada). After staming of the NP of P, the cells were wished and developed with fluorescent softhocymate compared goar anti-mouse immunoglobulin G or anti-rabbit immunoglobul a G-labeled with rhodamine. The samples were mounted with Mosaol 40-88 (Aldrich, Inc.) and analyzed by using a 40 ° or 63 ° objective and a licical CLSM confocal imaging

### RESULTS

The amino terminus of the influenza A virus NP contains a binding site for NPI-1 and NPI-3. In order to study the interaction of the viral NP and cellular NPI-1 and NPI-3 proteins, the yeast two-hybrid system was used (8, 11). The NP and its deletion mutants were expressed in yeast as LexA-NP fusion proteins. The plasmids expressing the NPI-1 or NPI-3 fusion proteins with the transcriptional activation domain had been obtained previously (20, 21). Plasmids containing the NP tu-

sion gene and the NPI-1 or NPI-3 fusion gene were cotransfected into yeast cells together with a reporter plasmid expressing β-galactosidase. Transformants were then transferred to X-Gal plates to allow easy screening for interactions between the NP mutants and either NPI-1 or NPI-3. Earlier results had suggested that a sequence associated with nuclear accumulation of the influenza A virus NP in Xenopus oocytes was located between amino acids 327 and 345 (5). Thus, we first made deletions in this region. However, binding activity of the resulting deletion mutants to either NPI-1 or NPI-3 was not affected in the yeast two-fusion protein assay (constructs A and B [Fig. 1a]). When we made deletions of the N-terminal 93 or 73 amino acids in constructs C and E, respectively, interaction with either of the NPI proteins was completely eliminated. Conversely, constructs D and F, which contained only the N-terminal 93 and 73 amino acids, respectively, bound to both

In order to examine the cellular localization of the mutant NP proteins, several constructs were cloned into an expression plasmid behind a cytomegalovirus (CMV) promoter. Transfection of these plasmids into HeLa cells was monitored 24 h later by indirect immunofluorescence (Fig. 1b). Mutants A and B, which lack the putative nuclear accumulation signal (5), localized to the nucleus, whereas construct E, which had an N-terminal deletion of 73 amino acids, was clearly cytoplasmic. These results suggested a correlation between NPI binding domains and an activity for nuclear localization in mammalian cells.

Amino acids 1 to 20 of the NP contain a binding site for NPI-I and NPI-3 and NLS activity. The following experiments were done to further define the N-terminal domain containing

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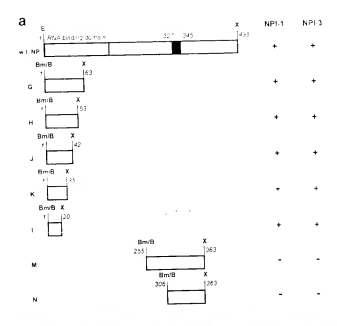
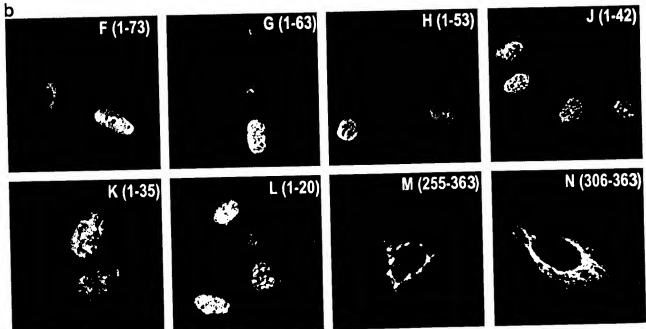


FIG. 2. Ammo acids I to 20 of the influenza A virus NP possess NI Sactivity. (g) Interactions of NP deletion mutants and NPI-1 and NPI-3 in the yeast two-hybrid system. The NP deletion mutants were closed into pI G202 and assayed as described in the legend to Fig. 1, Restriction enzyme sites used for subcoming of NP fragments are as follows: B. Bg/II. E. E. GRI, X. XI/ol. Bar. BamHI (8) Localization of tusion prot. inscontaining NP fragments and measles via P protein in HeLa cells. The fusion genes were cloned into PECE (provided by Eonald Harty), transfected into HeLa cells, and expressed as NP-P fusion proteins. The localization of the fusion proteins was analyzed by indirect immunofluorescence assay 24 h posttransfection, using anti-mearles virus P protein antibody PVI (kindly provided by Dalius J. Briedis). Constracts F to L (Fig. Ia and 2a) contain NP-specific amino acids as indicated. Constructs M and N contain NP-specific amino acids 255 to 363 and 306 to 363, which include the sequences previously thought to possess NLS activity (5). Constructs K. M. and N were photographed with the 63+ objective. The remaining photographs were taken with the 40+ objective.



		NPI-1 NPI-3	
w.t.	MASOGTKESTEÇMETDGERÇ	+	+
<b>A</b> 3	MAAQGTKRSYEUMETIGERT	-	+
A4	MASAGTKFSYROMETLGEF.	+	+
<b>A</b> 5	MASQATKRSYEQMETDGER. <sub>2</sub>	-	+
<b>A</b> 6	MASQGAKRSYEQMETEGERD	-	-
<b>A</b> 7	MASQ STARSYECMETEGER )	-	-
A8	MAS GTKASYECMITE GER D	-	-
<b>A</b> 9	MASQ GTKRAYEQMETEGER D	-	-
A10	MASIGTKRSAE METIGER.	-	+
A11	MASĮGTKRSY <b>A</b> ĮMETLGERĮ	+	+
A12	MASQGTKRSYE <b>A</b> METUGER D	+	+
A13	MAS DETKRISYET ARTDIGER .	-	-
A14	MAS I STEE SYE (MATIGER )	+	+
A15	MAS GTYPSYEGME <b>A</b> DGEF D	+	+
A16	MAS I GTHE SYEQMET <b>A</b> GEF D	+	+
A17	MAS (GTYLSYE (METE <b>A</b> EF )	+	+
A18	mas įgtkesye "meti. G <b>a</b> e į	+	+
A19	mas pstkrsyermetl se <b>a</b> q	+	+
A20	mas jetkisyet meti gel <b>a</b>	+	+
D5	MTKRSYEQMETUGERQ	-	+
D5A7	MTARSYEQMETDGEF 2	-	-
D5A8	MTK <b>A</b> SYEÇMETDGEH.	-	-
D5D9	MTKR-YE_METDGER_2	-	-
D5D10	MTHRS-EQMETIGERD	-	-
D5A19	MTKRSYE_METEGE <b>A</b> Q	-	+
NPI-1 Motif	a. amied averily		
NPI-3 Motif	TKRSADAM		
		N DE LANDE	A costore will

FIG. 3. Fine a solong and alarm a scanning of the NPI-1 NPI-3 interacting site on the NP. M. For and deficions of the first 20 annino acids of the NP were closed into p4 (120) and issued as described in the legend to Fig. 1 for binding to NPI-1 to to NPI-3. A rine mutations are in boldface and underfined 15 objects indicate a bettor. All numer NPI-1 and NPI-3 binding motifs are indicated at the bottom, wat, wild type

the NPI-1 NPI-3 binding site. Carboxy-terminal deletions were made in the first 73 amino acids, and fusions were constructed in vector pEG202. Lusions containing as few as 20 amino acids of the NP N terminus bind to both NPI-1 and NPI-3 when the yeast two-hybrid system is used (Fig. 2a). In order to test if these sequences function as NLS motifs, these N-terminal sethences were expressed as tusion proteins by using a reporter tein usually tound in the cytoplasm. Use of a reporter was necessary because the (mall N-terminal NP fragments could not be visualized with the available NP-specific antisera. Fusion constructs were made with the measles virus P protein. which localizes to the cytoplasm (12). Fusion constructs containing the 73 63, 53 42, 35, and 20 N terminal amino acids of the NP relocate the reporter protein into the nucleus (constructs F, G, H. J. L. and L. respectively [Fig. 2b]). Control constructs which express NP-specific amino acids 255 to 363 and 306 to 363 in the context of the measles virus P protein are and only in the cytopla in, confirming the absence of a nuir accumulation signal in these sequences

Thus, the NFI-1 NPI-3 binding site of the NP also functions

Fine mapping of the NPI-I/NPI-3 binding site. In order to identify the specific amino acids involved in binding to NPI-1 and NPI-3, we performed a mutational analysis, including alanine scanning mutagenesis, followed by assaying of the mutants in the yeast two-hybrid system. Figure 3 lists 24 mutants which were analyzed. The results suggest slight differences in the binding site recognized by NPI-1 and NPI-3. First, it appears that the five amino acids at the N terminus (or at least amino acids two to five) are not required for binding to NPI-3. This conclusion is based on the binding characteristics of mutants A3, A4, A5, D5, and D5A19. Second, mutant A10 is recognized by NPI-3 and not by NPI-1, again suggesting a difference in the binding pattern. These data reveal minimal binding motifs of SxGTKRSYxxM and TKRSxxxM for NPI-1 and NPI-3, respectively.

In addition, the number of amino acids between the serine and methionine is important for NPI-3 binding to the NP. Mutant D5 is recognized by NPI-3, but D5D10 is not. Since mutatton of Y10 to alanine does not disrupt the NPI-3 binding site (mutant A10), but deletion of this residue does ablate binding, we conclude that a three-amino-acid spacer between the serine and methionine is necessary for recognition of this NLS by NPI-3.

Several mutants were also cloned as fusion genes (with the measles virus P gene) into the CMV-driven expression plasmids. They were then transfected into HeLa cells and analyzed for cellular localization by indirect immunofluorescence (Fig. 4). Mutants A7, A8, D5A7, D5A8, and D5D10, which have lost the ability to bin-leither NPI-1 or NPI-3, localize to the cytoplasm. However, mutants D5A19 and D5, which retain NPI-3 binding activity, are clearly nuclear in this assay. These results suggest a good correlation between NPI binding and NLS activity. It also appears that the binding site for NPI-1 and NPI-3 is contained within the first 13 amino acids at the N terminus. Finally, the data suggest a redundancy of two slightly different NLS signals mediated by binding either to NPI-1 or to NPI-3.

Presence of an NLS whose activity is not mediated by binding to NPI-1 or NPI-3. In order to confirm the localization of the NPI-1 NPI-3 binding site in the full-length NP molecule, the first 12 amino acids were deleted (construct D12 [Fig. 5a]). As expected, this mutant did not bind NPI-1 or NPI-3 in the yeast two-hybrid system. However, this construct-when inserted into the CMV-driven expression plasmid and transfected into HeI a cells--unexpectedly localized mostly to the nucleus (Fig. 5b). This phenomenon was further investigated by examining a construct that lacked amino acids 1 to 12 and 255 to 340. Although this mutant lacked the NPI binding site and the putative NLS previously described (5), it again was found predominantly in the nucleus. We then made an attempt to further define the domain which is associated with this NLS activity. Construct O contained the D5A7 mutation at the N terminus and was truncated at amino acid position 254 (so that it also lacked an NPI binding site). This construct localized in the cytoplasm when expressed as a fusion protein (containing the measles virus P protein) (Fig. 5b). We then discovered that

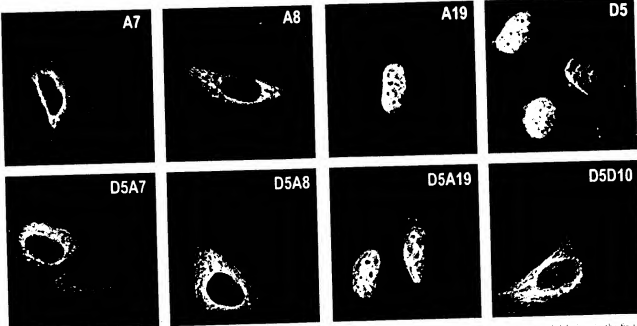


FIG. 4. Mutations and deletions in the first 20 amino acids of the influenza A virus NP fragment affect NLS activity. The mutations and deletions in the first 20 amino acids of the NP were cloned into the fusion gene vector PECE, transfected into HeLa cells, and expressed as fusion proteins (containing the measles virus P protein). The localization of the fusion proteins was analyzed by indirect immunofluorescence assay 24 h posttransfection, using anti-measles virus P protein antibody PVI.

constructs O and E (neither of which alone carries an NLS) allowed localization to the nucleus when cotransfected. This result suggests that the NLS activity associated with the D12 mutant is mediated by noncontiguous regions in the NP molecule and that NP deletions O and E can interact in *trans*, thereby reconstituting a conformational NLS.

## DISCUSSION

Nuclear protein import is accomplished by two sequential events: docking at and translocation through the nuclear membrane (9, 10, 16–19, 22). Through the use of the yeast two-hybrid system, with the NP as "bait," we identified two proteins, NPI-1 and NPI-3 (docking proteins), which were shown to be involved in the nuclear import of the influenza virus NP and of viral RNA (20, 21). The present study was done in order to identify the NPI binding site on the NP and to study its biological activity. We assayed the interaction of the NP with NPI-1 and NPI-3 in the yeast two-hybrid system by using NP deletion mutants and found the NPI binding site localized to the amino terminus (Fig. 1). Furthermore, the amino terminus of the NP showed NLS activity, since it could confer nuclear localization on a protein (the measles virus P protein) which normally localizes to the cytoplasm (Fig. 2).

In contrast, earlier studies (5) had suggested that nuclear accumulation of the NP was dependent not on the N terminus but rather on amino acids 327 to 345 of the protein. The proposed nucleophilic signal (AAFEDLRVLS) was found to be important for the nuclear accumulation of the NP as well as of NP-globin fusion proteins in *Xenopus* oocytes. However, we found that expression of NP mutants A and B, lacking amino acids 255 to 356 and 255 to 340, respectively, showed nuclear localization (Fig. 1b). Furthermore, two fusion proteins containing amino acids 255 to 363 and 306 to 363 (both encompassing the putative NLS) of the viral NP were cytoplasmic in HeLa cells (Fig. 2b). These findings suggest that the earlier data concerning the nucleophilic signal of the NP (located

between amino acids 327 and 345) were most likely an artifact of the *Xenopus* system and could not be verified in mammalian cells (at least not in HeLa cells).

A more refined analysis of the NPI binding site suggested that the N-terminal 13 residues of the NP contain the necessary amino acids involved in binding of NPI-1 and NPI-3. Alanine scanning identified the motif TKRSxxxM as sufficient for NPI-3 binding as well as for NLS activity. Mutants which have lost the ability to bind NPI-1 (but retain NPI-3 binding activity) are able to direct a fusion protein to the nucleus and thus appear to retain NLS activity by virtue of their interaction with NPI-3 (mutants D5 and D5A19 [Fig. 4]). In a fractionated nuclear import assay either NPI-1 or NPI-3 was able to direct free NP and NP in the form of an RNP complex to the nucleus (21). Thus, there is a redundant mechanism in terms of the NLS located at the N terminus. Binding of the NP to either NPI-1 or to NPI-3 may facilitate docking to the nuclear membrane and subsequent translocation through the membrane.

Nothing is known about the differential expression of the docking proteins in various cell types, and thus it is not possible to identify the NPI(s) involved in natural influenza virus infections. It should be noted, however, that there appear to be additional members of the NPI-1-NPI-3 protein family. There are more than 50 human expressed sequence tags in the database at the National Center for Biotechnology Information. Sequence alignments of some of these expressed sequence tags with the region spanning nucleotides 800 to 1200 of NPI-I demonstrate that there are at least four different human NPI-1 protein family members (including NPI-1 and NPI-3). The sequence of a partial clone homologous to NPI-1 and NPI-3 was published by Adams et al. (1). It is thus possible that there is a high degree of redundancy (as well as specificity) for the nuclear import of NLS-containing proteins and that the nuclear localization of proteins may be mediated by the presence of different NPI proteins in the cell.

The NLS motifs SxGTKRSYxxM and TKRSxxxM, which were defined by the interaction with NPI-1 and NPI-3, respec-

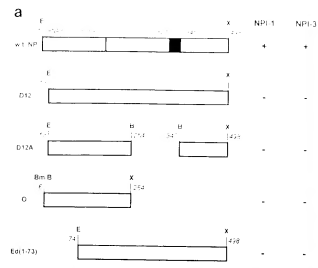
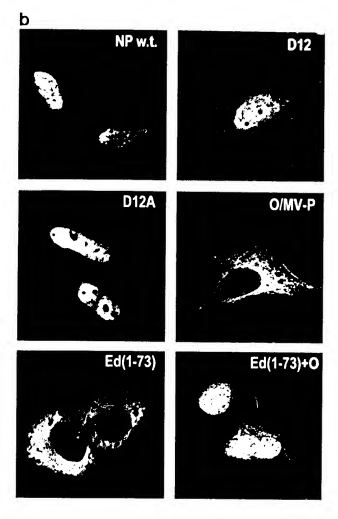


FIG. 5. NLS activities of NP mutants lacking an NPI binding site. (a) NP mutants were constructed and cloud into ph/G202 (see Materials and Methods) and user ed as described in the legend to Fig. 1. Restriction enzyme sites are indicated as follows: 1. Jan/RL B. Bg/IL X. Alic 1 (b) Localization of NP mutants Helia cells. NP mutants D12. D12A, and L were cloned into pcDNAI Amp and transfected into Helia cells. The O'MV-P fusion gene consists of the NP tragment O gene and the measles virus P gene. NP tragment O contains mutation D5A7 (deletion of five amino acids and change to alamne at position 7 (a) the N terminus and is truncated at amino acid position 254 at the C terminus. The localization of fusion proteins was analyzed by indirect immunofluorescence assay 24 h postriansfection, using the following anti-NP antibodies polyclonal anti-N-31 antibody for staining of constructs E d(1-73) and D12A and for the cotransfection of constructs F d(1-73) and D12 constructs, and anti-meades virus P protein antibody PV1 for staining of the O MV-P tusion protein. Ed(1-73) +O, cotransfection of constructs E d(1-73) and O.

tively, do not resemble those identified in the similar virus 40 T antigen and nucleoplasmin (6). In addition to these two classical NLS sequences, nonconventional NLS motifs have been described (24–26). None of these appear to be similar or identical to those identified in the influenza virus NP.

RNA binding domains of the NP have been mapped to amino acids 1 to 77, 79 to 188, and 91 to 188 (1a, 14). However, at this time RNA binding domains have not been fine mapped on the NP, so we do not know whether the NPI binding sites overlap the RNA binding domain. Import of the viral RNA most likely involves a piggyback mechanism by which the RNA is bound to viral NP that gets transported into the nucleus (21).

Unexpectedly, we also identified an NLS activity which does Pot appear to be mediated by binding to either NPI-1 or NPI-3 (constructs D12 and D12A [Fig. 5b]). This NLS activity is most bkely the result of a discontinuous amino acid sequence on the MP, since cotransfection of two fragments (constructs O and E [Fig. 5b]) results in nuclear localization measured by immunofluorescence. The signals have not been further investigated, and the mechanism by which this nuclear transport is effected is not known. Since each fragment alone has a cytoplasmic localization, the two fragments may interact directly or indirectly and thereby reconstitute a biologically active NLS. Indeed, data from the two-hybrid assay indicate that full-length SP can interact with itself (data not shown). In any case, this NLS activity does not appear to involve binding to NPI-1 or NPI-3 (as measured in the yeast two-hybrid system). This signal may be weaker than that associated with the N terminus, since the constructs containing the bipartite NLS show partial cytoplasmic localization (mutant D12 [Fig. 5b]). However, we



cannot be certain about the relative strengths of these NLS activities, since we lack a quantitative in vivo assay. This bipartite NLS may also represent an NLS which is assembled from adjacent NPs in RNP complexes rather than a signal assembled from domains normally contained within a single NP molecule. Finally, based on the analysis of the amino acid sequence of NP, the above-described NLS activity does not appear to be mediated by classical NLS sequences.

In summary, the present data characterize the NPI binding site on the influenza A virus NP. The N-terminal 13 amino acids contain a motif which is associated with binding to NPI-1 and NPI-3. In addition, this sequence possesses an NLS activity, as can be demonstrated by the nuclear localization of fusion proteins which contain this short motif. The N-terminal 13 amino acids of influenza A virus NPs are highly conserved. Of the 56 avian and mammalian NP sequences available in the data bank, 54 are identical to the sequence of the A PR 8.34 virus NP. The specific NPI binding motifs identified in the influenza A virus NP appear not to be present in influenza B and C virus NPs. It will be important to examine whether the NPs of these viruses also bind to NPI-1 and NPI-3 and whether these binding sites have NLS activities

### ACKNOWLEDGMENTS.

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ship at diagrams) the meast is virus P protein and monoclored amploals duveled arabist the influenza virus NP protein, it specified.

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